510(k) Summary

per 21 CFR §807.92

Submitter's Name and Address Boston Scientific Corporation

One Scimed Place Maple Grove, MN 55311

Contact Name

Carol Tiffany

Senior Regulatory Affairs Specialist

and Information

Phone: 763-494-1106 Fax: 763-494-2222

Fax: 763-494-2222 e-mail: carol.tiffany@bsci.com

Date Prepared

29 April 2014

Proprietary Name Sterling™ Over-the-Wire PTA Balloon Dilatation Catheter

Common Name

Percutaneous Catheter

Product Code

LIT - Catheter, Angioplasty, Peripheral, Transluminal

Classification

Class II, 21 CFR Part 870.1250 - Percutaneous Catheter

Predicate Device(s) Sterling OTW PTA Balloon Dilatation Catheter K132430 October 17, 2013

Device Description The Sterling OTW Percutaneous Transluminal Angioplasty (PTA) Balloon Dilatation Catheter is a high performance balloon catheter for peripheral indications. The device features an ultra-low profile, semi-compliant balloon combined with a low profile tip. The catheter is compatible with either 0.014 in (0.36 mm) or 0.018 in (0.46 mm) guidewires.

The Sterling OTW PTA Balloon Dilatation Catheter is an Over-The-Wire (OTW) catheter with a semi-compliant balloon fixed at the distal tip. The balloon catheter has a coaxial shaft design. The outer lumen is used for inflation of the balloon, and the wire lumen permits the use of guide wires 0.014 in or 0.018 in to facilitate advancement of the catheter to and through the stenosis to be dilated. The balloon is designed to provide an inflatable segment of known diameter and length at recommended pressures. The catheter includes a tapered tip to facilitate advancement of the catheter to and through the stenosis. Two radiopaque marker bands (one proximal and one distal), in conjunction with fluoroscopy, enable accurate positioning of the balloon.

The effective lengths of the balloon catheter are 90 cm and 150 cm. Markers on the 90 cm effective length catheter indicate the exit of the dilatation catheter tip out of the guiding catheter (one at 50 cm and two at 60 cm). Markers on the 150 cm effective length catheter indicate the exit of the dilatation catheter tip out of the guiding catheter (one at 90 cm and two at 100 cm). The proximal portion of the catheter includes one female Luerlock port connected to the inflation lumen, and one female Luerlock port for guidewire lumen.

The balloon lengths are available in 20, 30, 40, 60, 200 and 220 mm sizes with diameters of 1.5, 2.0, 2.5, 3.0, 3.5 and 4.0 mm. The combinations of balloon sizes are displayed below.

Balloon (mm)	Balloon Length (mm)						
	20	30	40	60	200	220	
1.5	90cm 150cm	,	90cm 150cm				
2.0	90cm 150cm	90cm 150cm	90cm 150cm	90cm 150cm		90cm 150cm	
2.5	90cm 150cm	90cm 150cm	90cm 150cm	90cm 150cm		90cm 150cm	
3.0	90cm 150cm	90cm 150cm	90cm 150cm	90cm 150cm		.90cm 150cm	
3.5	90cm 150cm	90cm 150cm	90cm 150cm	90cm # 150cm		90cm 150cm	
4.0		,			90cm 150cm	90cm 150cm	

Intended Use/ Indications for Use of Device The Sterling OTW PTA Balloon Dilatation Catheter is indicated for Percutaneous Transluminal Angioplasty (PTA) in the peripheral vasculature, including iliac, femoral, popliteal, infra-popliteal, and renal arteries, and for the treatment of obstructive lesions of native or synthetic arteriovenous fistulae. This device is also indicated for post-dilatation of balloon expandable and self-expanding stents in the peripheral vasculature.

Comparison of Technological Characteristics

The Sterling OTW catheter will incorporate a substantially equivalent design, packaging, fundamental technology, manufacturing, materials, sterilization and intended use as those featured in the predicate BSC Sterling OTW Balloon Dilatation Catheter.

Characteristic	Comparison to Sterling OTW Predicate		
Manifold .	Same material. Same design serving the same function.		
Strain Relief	Same material. Same design serving the same function.		
Inner Shaft/Outer Shaft	Same material. Same design serving the same function.		
Bailoon	Same material. Same design serving the same function.		
Marker Bands	Same component. Same design serving the same function.		
Proximal Marks	Same material. Same design serving the same function.		
Coating	Same material. Same design serving the same function.		
Bumper Tip	Same material. Same design serving the same function.		
Sterilization Method/SAL	Same method and same level of assurance.		
Balloon Diameters	Smaller diameters than the predicate range.		
Balloon Lengths	200 and 220 mm are the same as predicate range. Adding balloon lengths of 20, 30, 40, and 60 mm.		
Usable Catheter Lengths	Same lengths.		
Rated Burst Pressure (RBP)	Same Rated Burst Pressure.		
Sheath/Guide Compatibility	Compatibility the same for larger sizes, slightly different compatibility depending on balloon diameters.		
Packaging	Same function and design.		
Guidewire	Same Compatibility.		
Manufacturing	Manufactured on the same manufacturing lines.		

Performance Data

Bench testing was performed to support a determination of substantial equivalence. The results of these tests provide reasonable assurance that the proposed device has been designed and tested to assure conformance to the requirements for its intended use. No new safety or performance issues were raised during the testing and, therefore, these devices may be considered substantially equivalent to the predicate devices.

The following bench testing was completed on the Sterling™ OTW PTA Balloon Dilatation Catheter:

Bench

Balloon Compliance

Balloon Nominal Diameter

Balloon Rated Burst Pressure (RBP)

Burst in a Stent

Balloon Multiple Inflation

Balloon Body Length

Crossing Profile

Guidewire Movement

Full Catheter Tensile Extension

Sheath Withdrawal

and Deflation

Balloon Multiple Inflation

Marker Band to Balloon

in a Stent

Alignment

Proximal Balloon Bond and Shaft Tensile Strength

Deflation Time

Conclusion

Based on the Indications for Use, technological characteristics, safety and performance testing, the Sterling OTW PTA Balloon Dilatation Catheter has been shown to be appropriate for its intended use and is considered to be substantially equivalent to the Sterling OTW PTA Balloon Dilatation Catheter (K0132430 cleared October 17, 2013).



Food and Drug Administration 10903 New Hampshire Avenue Document Control Center – WO66-G609 Silver Spring, MD 20993-0002

July 16, 2014

Boston Scientific
Ms. Carol Tiffany
Regulatory Affairs Specialist
One Scimed Place
Maple Grove, MN 55311-1566

Re: K141112

Trade/Device Name: Sterling Over-The Wire PTA Balloon Dilatation Catheter

Regulation Number: 21 CFR 870.1250 Regulation Name: Percutaneous Catheter

Regulatory Class: Class II

Product Code: LIT
Dated: June 13, 2014
Received: June 16, 2014

Dear Ms. Tiffany:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803); good manufacturing practice requirements as set

forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to

http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm.

Sincerely yours,

Kenneth J. Cavanaugh -S

for

Bram D. Zuckerman, M.D.
Director
Division of Cardiovascular Devices
Office of Device Evaluation
Center for Devices and Radiological Health

Enclosure

Indications for Use

510(k) Number (if known):							
Device Name:	Sterling™ Over-	the-Wire (OTW) F	PTA Balloon Dilatation Catheter					
Indications for U	se:							
Transluminal An popliteal, infra-ponative or synthetic	gioplasty (PTA) i opliteal, and rena tic arteriovenous	n the peripheral va I arteries, and for dialysis fistulae. T	is indicated for Percutaneous asculature, including iliac, femoral, the treatment of obstructive lesions of his device is also indicated for postgrents in the peripheral vasculature.					
Prescription Use _ (Part 21 CFR 801		AND/OR	Over-The-Counter Use (21 CFR 801 Subpart C)					
(PLEASE DO NOT WRITE BEŁOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED)								
(Concurrence of C	DRH, Office of De	vice Evaluation (ODE)					

